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FILE CONTENT:1840 - 10 May 2008 VOL 148 ISS 20

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           CA reference information (SO, PY). (Default)
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            Summary for all hit reactions and fields containing
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## L3 ANSWER 1 OF 10 CASREACT COPYRIGHT 2008 ACS on STN

81%

REF: Heterocycles, 71(1), 39-48; 2007 NOTE: key intermediate CON: STAGE(1) 1 hour, reflux; 2 hours, reflux STAGE(2) room temperature -> 130 deg C; 3 hours, 130 deg C

=> d cbib abs crd 1-10

146:421938 One-pot conversion of 2-nitrobenzonitriles to quinazolin-4(3H)-ones and synthesis of gefitinib and erlotinib hydrochloride. Chandregowda, Venkateshappa; Rao, Gudapati Venkateswara; Reddy, Goukanapalli Chandrasekara (Vittal Mallya Scientific Research Foundation, Bangalore, 550004, India). Heterocycles, 71(1), 39-48 (English) 2007. CODEN: HTCYAM. ISSN: 0385-5414. Publisher: Japan Institute of Heterocyclic Chemistry.

AB A simple and efficient one-pot conversion of 2-nitrobenzonitriles to quinazolin-4(3H)-ones involving reduction, formylation, hydrolysis and cyclization is reported. E.g., quinazolin-4(3H)-one derivative I was prepared with 85% yield by reacting the corresponding 2-nitrobenzonitrile II with hydrazine using FeCl3 in MeOH/H2D followed by treating the reaction mixture with formic acid and HCl. These quinazolinones have been used for making in economical way the anticancer drug mols. gefitinib (Iressa) and erlotinib HCl (Tarceva).

NOTE: key intermediate

CON: STAGE(1) 1 hour, reflux; 2 hours, reflux

STAGE(2) room temperature -> 130 deg C; 3 hours, 130 deg C

```
RX(37) OF 57 - 2 STEPS
```

O-CH2-CH2-OMe

1.1. HNO3, Water,

Ac20 1.2. NH4OH, Water 2.1. FeCl3, N2H4,

Water, MeOH 2.2. HCO2H, HCl, Water

> MeO-CH2-CH2-O MeO-CH2-CH2-C

NOTE: 1) 75% overall yield from 3,4-dihydroxybenzaldehyde, regioselective, 2) key intermediate
CON: STEP(1.1) 45 - 50 deg C; 8 hours, 45 - 50 deg C
STEP(1.2) pH 8
STEP(2.1) 1 hour, reflux; 2 hours, reflux

STEP(2.2) room temperature -> 130 deg C; 3 hours, 130 deg C

RX(48) OF 57 - 3 STEPS

O-CH2-CH2-OMe

81%

NOTE: 2) 75% overall yield from 3,4-dihydroxybenzaldehyde,

regioselective, 3) key intermediate CON:

STEP(1.1) 25 deg C -> reflux; 1 hour, reflux STEP(1.2) room temperature -> 110 deg C; 3 hours, 110 deg C

STEP(1.3) pH 8 STEP(2.1) 45 - 50 deg C; 8 hours, 45 - 50 deg C STEP(2.2) pH 8

STEP(3.1) 1 hour, reflux; 2 hours, reflux

STEP(3.2) room temperature -> 130 deg C; 3 hours, 130 deg C

RX(49) OF 57 - 4 STEPS

81%

NOTE: 3) 75% overall yield from 3,4-dihydroxybenzaldehyde,

regioselective, 4) key intermediate
STEP(1.1) room temperature -> 100 deg C; 3 hours, 100 deg C CON: STEP(2.1) 25 deg C -> reflux; 1 hour, reflux

STEP(2.2) room temperature -> 110 deg C; 3 hours, 110 deg C

STEP(2.3) pH 8

STEP(3.1) 45 - 50 deg C; 8 hours, 45 - 50 deg C STEP(3.2) pH 8

STEP(4.1) 1 hour, reflux; 2 hours, reflux

STEP(4.2) room temperature -> 130 deg C; 3 hours, 130 deg C

- ANSWER 2 OF 10 CASREACT COPYRIGHT 2008 ACS on STN 146:274315 Improved synthesis of substituted 6.7-dihydroxy-4
  - quinazolineamines: tandutinib, erlotinib and gefitinib. Knesl, Petr; Roeseling, Dirk; Jordis, Ulrich (Institute of Applied Synthetic Chemistry, Vienna University of Technology, Vienna, 1060, Austria). Molecules, 11(4), 286-297 (English) 2006. CODEN: MOLEFW. ISSN: 1420-3049. URL: http://www.mdpi.org/molecules/papers/11040286.pdf Publisher: Molecular Diversity Preservation International.
- AB The synthesis of three substituted 6,7-dihydroxy-4-quinazolineamines: tandutinib, erlotinib and gefitinib in improved yields is reported. The intermediates were characterized by NMR and the purities determined by HPLC.

RX(12) OF 81

$$\begin{array}{c} \text{MeO-CH}_2\text{-CH}_2\text{-O} \\ \text{MeO-CH}_2\text{-CH}_2\text{-O} \\ \text{NH}_2 \end{array} \qquad \begin{array}{c} \text{Formamide} \\ \text{NH}_2 \\ \end{array}$$

84%

CON: 12 hours, 165 - 170 deg C

1. PtO2, H2, MeOH 2. Formamide

NO<sub>2</sub> MeO-CH2-CH2-O

84%

CON: STEP(1) room temperature, 50 psi STEP(2) 12 hours, 165 - 170 deg C

RX(53) OF 81 - 3 STEPS

1. HNO3, Water, AcOH 2. PtO2, H2, MeOH

MeO-CH2-CH2-O MeO-CH2-CH2-0 3. Formamide

84%

CON: STEP(1.1) 30 minutes, 0 - 5 deg C; 24 hours, room temperature STEP(2) room temperature, 50 psi STEP(3) 12 hours, 165 - 170 deg C

RX(54) OF 81 - 4 STEPS

NOTE: 1) alternative preparation shown CON: STEP(1.1) 20 minutes, room temperature

STEP(1.1) 5 days, reflux STEP(2.1) 30 minutes, 0 - 5 deg C; 24 hours, room temperature

STEP(3) room temperature, 50 psi STEP(4) 12 hours, 165 - 170 deg C

- L3 ANSWER 3 OF 10 CASREACT COPYRIGHT 2008 ACS on STN
- 145:271722 Fluorine-18 labeling of 6,7-disubstituted anilinoquinazoline derivatives for positron emission tomography (PET) imaging of tyrosine kinase receptore: Synthesis of 18F-Iressa and related molecular probes. Seimbille, Yann; Phelps, Michael E.; Czernin, Johannes; Silverman, Daniel H. S. (Ahmanson Biological Imaging Division, Department of Molecular and Medical Pharmacology, University of California, Los Angeles, CA, 90095-6942, USA). Journal of Labelled Compounds & Radiopharmaceuticals, 48(11), 829-843 (English) 2005. CODEN: JLCRD4. ISSN: 0362-4803. Publisher: John Wiley & Sons Ltd..
- AB Inhibitors of tyrosine kinase enzymic activity are a promising new class of antineoplastic agents. Although clin. studies performed over the last decade give more insight on the potential therapeutic applications of such drugs, identification of the individual patients who might benefit from them remains a major challenge. The authors have developed a synthetic strategy for the production of a wide variety of radiolabeled 6,7-disubstituted 4-anilinequinazolines suitable for non-invasive imaging of tyrosine kinase receptors to predict therapy effectiveness. Three new F-18 labeled radiopharmaceuticals based on the therapeutic agents Tarceva, Iressa, and 2D647% were synthesized. Decay-corrected yields were 25-40% for a total synthesis time of 120 min, thus providing F-18 labeled tyrosine kinase inhibitors in quantities and times practical for use as PET radiopharmaceuticals.

RX(23) OF 163

Formamide, Ammonium formate

NOTE: Niementowski condensation CON: 3 hours, 160 deg C

RX(45) OF 163 - 2 STEPS

- 1. Pd, H2, MeOH
- 2. Formamide,
  Ammonium formate

NOTE: 2) Niementowski condensation CON: STEP(1) room temperature STEP(2) 3 hours, 160 deg C RX(74) OF 163 - 3 STEPS

NOTE: 3) Niementowski condensation CON: STEP(1.1) 15 minutes, 0 - 5 deg C; 2 hours, room temperature STEP(2) room temperature

STEP(3) 3 hours, 160 deg C

RX(75) OF 163 - 4 STEPS

NOTE: 4) Niementowski condensation CON: STEP(1) 72 hours, reflux

STEP(2.1) 15 minutes, 0 - 5 deg C; 2 hours, room temperature STEP(3) room temperature

STEP(4) 3 hours, 160 deg C

L3 ANSWER 4 OF 10 CASREACT COPYRIGHT 2008 ACS on STN

145:249220 Preparation of 4-(indol-3-yl)quinazolines as epidermal growth factor receptor inhibitors. Loewe, Werner; Lueth, Anja (Freie Universitaet Berlin, Germany). PCT Int. Appl. WO 2006084882 A2 20060817, 23pp. DESIGNATED STATES: W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BW, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, EG, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KM, KN, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, LY, MA, MD, MG, MK, MN, MW, MX, MZ, NA, NG, NI, NO, NZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, SM, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA; RW: AT,

BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, IS, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG, TR. (German). CODEN: PIXXD2. APPLICATION: WO 2006-EP50813 20060209. PRIORITY: DE 2005-102005007151 20050211.

AB Title compds. I [Rl = methoxy, ethoxy, propoxy, etc.; R2 = H, methoxy, etc.; R3 = (R3')n; R3' = halo, halo substituted benzyloxyl, alkyl, etc.; n = 1-3] and their pharmaceutically acceptable salts were prepared For example, Mg mediated coupling of 4-chloro-6, 7-dimethoxyquinazoline and 5-bromoindole afforded quinazoline II. In epidermal growth factor receptor inhibition assays, 4-examples of compds. I exhibited 61-81% inhibition at 0.1 µM.

## RX(5) OF 18

CON: 24 hours, reflux

- L3 ANSWER 5 OF 10 CASREACT COPYRIGHT 2008 ACS on STN
- 144:370039 Synthesis and biological evaluation of allenic quinazolines using palladium-catalyzed hydride-transfer reaction. Nakamura, Hiroyuki; Onagi, Shinya (Department of Chemistry, Faculty of Science, Gakushuin University, Mejiro, Tokyo, 171-8588, Japan). Tetrahedron Letters, 47(15), 2539-2542 (English) 2006. CODEN: TELEAY. ISSN: 0040-4039. Publisher: Elsevier B.V..
- AB Allenic quinazolines were designed as mimics of Tarceva, which is an epidermal growth factor receptor (EGFR) tyrosine kinase inhibitor, and synthesized from the corresponding 4-(iodoanilino)quinazolines or 4-(iodophenoxy)quinazolines with N,N-dicyclohexylprop-2-ynylamine by the Sonogashira coupling followed by palladium-catalyzed hydride-transfer

reaction. Cell growth inhibition of the new compds. toward A431, Kato III, SKBR3, and HepG2 was examined One of the compds. synthesized, showed a similar cell growth inhibition to Tarceva. Moreover, two other compds. exhibited a specific growth inhibition toward Kato III cells (IC50 = 12 and 4.7  $\mu\text{M}$ , resp.), although a significant inhibition toward other three cell lines was not observed at a 100  $\mu\text{M}$  concentration of compds.

RX(16) OF 171

Formamide\_

CON: 160 deg C

MeO-CH<sub>2</sub>-CH<sub>2</sub>-O

61%

CON: reflux

RX(21) OF 171 MeO-CH<sub>2</sub>-CH<sub>2</sub>-O N + K2CO3, Me2CHOH OH MeO-CH<sub>2</sub>-CH<sub>2</sub>-O 
$$\times$$
 N +  $\times$  MeO-CH<sub>2</sub>-CH<sub>2</sub>-O  $\times$  MeO-CH<sub>2</sub>-CH<sub>2</sub>-O  $\times$  MeO-CH<sub>2</sub>-CH<sub>2</sub>-O

CON: reflux

CON: 60 deg C

Pd(PPh3)4, CuI, Et3N, MeCN

CON: 60 deg C

CON: 100 deg C

Pd complex, R:76858-94-1, CHC13

$$\begin{array}{c} \mathsf{MeO-CH_2-CH_2-O} \\ \mathsf{MeO-CH_2-CH_2-O} \\ \mathsf{O} \\ \mathsf{CH-C-CH_2} \end{array}$$

61%

CON: 100 deg C

1. Pd, H2, EtOH 2. Formamide

CON: STEP(2) 160 deg C

61%

99%

CON: STEP(1) reflux STEP(2) reflux

CON: STEP(1) reflux STEP(2) reflux

CON: STEP(1) reflux STEP(2) 60 deg C

RX(60) OF 171 - 2 STEPS MeO-CH<sub>2</sub>-CH<sub>2</sub>-O N + 
$$\frac{CH_2-C=CH}{CH}$$
 (Step 2)

1. K2CO3, Me2CHOH

2. Pd(PPh3)4, CuI,
Et3N, MeCN

CON: STEP(1) reflux STEP(2) 60 deg C

RX(63) OF 171 - 2 STEPS 
$$\label{eq:meo-ch2-ch2-ch2-o} \mbox{MeO-CH}_2\mbox{-C$$

CON: STEP(1) 60 deg C STEP(2) 100 deg C

CON: STEP(1) 60 deg C STEP(2) 100 deg C

RX(89) OF 171 - 3 STEPS

1. HNO3, Water, Ac20 2. Pd, H2, EtOH 3. Formamide

MeO-CH<sub>2</sub>-CH<sub>2</sub>-O 3. Formami

MeO-CH2-CH2-O

CON: STEP(3) 160 deg C

RX(90) OF 171 - 4 STEPS

CON: STEP(1) reflux STEP(4) 160 deg C

Formamide
 EtN(Pr-i)2, POC13,

PhMe 3. K2CO3, Me2CHOH

61%

CON: STEP(1) 160 deg C STEP(2) reflux STEP(3) reflux

99%

1. Pd, H2, EtOH

2. Formamide
3. EtN(Pr-i)2, POC13,
PhMe
4. K2CO3, Me2CHOH

61%

CON: STEP(2) 160 deg C STEP(3) reflux STEP(4) reflux RX(100) OF 171 - 4 STEPS

4. K2CO3, Me2CHOH

99%

RX(103) OF 171 - 3 STEPS

2. K2CO3, Me2CHOH 3. Pd(PPh3)4, CuI,

Et3N, MeCN

PhMe

24%

CON: STEP(1) reflux STEP(2) reflux STEP(3) 60 deg C

- 1. EtN(Pr-i)2, POC13, PhMe
- 2. K2CO3, Me2CHOH 3. Pd(PPh3)4, CuI, Et3N, MeCN

CON: STEP(1) reflux STEP(2) reflux

STEP(3) 60 deg C

RX(107) OF 171 - 4 STEPS

СН2-С≡СН

60%

- 1. Formamide 2. EtN(Pr-i)2, POC13, PhMe
- 3. K2CO3, Me2CHOH 4. Pd(PPh3)4, CuI, Et3N, MeCN

RX(108) OF 171 - 4 STEPS

CON: STEP(1) reflux STEP(2) 60 deg C STEP(3) 100 deg C

61%

CON: STEP(1) reflux STEP(2) 60 deg C STEP(3) 100 deg C RX(116) OF 171 - 4 STEPS

1. EtN(Pr-i)2, POC13,

- PhMe 2. K2CO3, Me2CHOH
- 3. Pd (PPh3) 4, CuI, Et3N, MeCN
- Pd complex, R:76858-94-1, CHC13

61%

RX(121) OF 171 - 5 STEPS

CON: STEP(1) reflux STEP(2) reflux STEP(5) 160 deg C RX(126) OF 171 - 5 STEPS

MeO-CH<sub>2</sub>-CH<sub>2</sub>-O N

1. HNO3, Water, Ac20
2. Pd, H2, EtoH
3. Formamide
4. Etn(Fr-1)2, POC13, PhMe
5. K2CO3, Me2CHOH

CON: STEP(3) 160 deg C STEP(4) reflux STEP(5) reflux

PhMe 5. K2CO3, Me2CHOH

99%

61%

CON: STEP(3) 160 deg C STEP(4) reflux STEP(5) reflux RX(130) OF 171 - 6 STEPS

RX(131) OF 171 - 6 STEPS

RX(134) OF 171 - 7 STEPS

1. MeOH

2. MeOCH2CH2Br

5. Formamide

RX(135) OF 171 - 7 STEPS

1. MeOH

2. MeOCH2CH2Br

5. Formamide

Pd, H2, EtOH
 Formamide
 EtN(Pr-i)2, POC13,

PhMe 4. K2CO3, Me2CHOH

5. Pd(PPh3)4, CuI, Et3N, MeCN

CON: STEP(2) 160 deg C STEP(3) reflux STEP(4) reflux STEP(5) 60 deg C

24%

CON: STEP(3) 160 deg C STEP(4) reflux STEP(5) reflux STEP(6) 60 deg C

3. Formamide

CON: STEP(3) 160 deg C STEP(4) reflux STEP(5) reflux STEP(6) 60 deg C (step 6)

RX(146) OF 171 - 7 STEPS

24%

CON: STEP(1) reflux STEP(4) 160 deg C STEP(5) reflux STEP(6) reflux STEP(7) 60 deg C

CON: STEP(1) reflux STEP(4) 160 deg C STEP(5) reflux STEP(6) reflux STEP(7) 60 deg C RX(150) OF 171 - 8 STEPS

RX(154) OF 171 - 5 STEPS 
$$\begin{array}{c} \text{MeO-CH}_2\text{-CH}_2\text{--}O \\ \text{MeO-CH}_2\text{--}\text{CH}_2\text{--}O \\ \end{array}$$
 
$$\begin{array}{c} \text{H}_2\text{C}\text{--}\text{C}\text{--}\text{C}\text{H} \\ \\ \text{CON: STEP(1) 160 deg C} \\ \text{STEP(2) reflux} \end{array}$$

CON: STEP(1) 160 deg C STEP(2) reflux STEP(3) reflux STEP(4) 60 deg C STEP(5) 100 deg C

CH<sub>2</sub>-C=CH

3.
4.
(step 4)

1. Formamide

2. EtN(Pr-i)2, POC13,

PhMe

3. K2CO3, Me2CHOH
4. Pd(PPh3)4, CuI,
Et3N, MeCN

5. Pd complex, R:76858-94-1, CHC13

RX(159) OF 171 - 6 STEPS 
$$\begin{array}{c} \text{MeO-CH}_2\text{--CH}_2\text{--O} \\ \text{MeO-CH}_2\text{--CH}_2\text{--O} \\ \text{N} \\ \text{CH---C---CH}_2 \end{array}$$

CON: STEP(2) 160 deg C STEP(3) reflux STEP(4) reflux STEP(5) 60 deg C STEP(6) 100 deg C

H<sub>2</sub>C — С — С н 55%

RX(167) OF 171 - 8 STEPS

HO OH (step 6) 
$$CH_2-CECH$$

$$CH_2-CECH$$

$$CH_2-CECH$$

$$(step 7)$$

$$MeO-CH_2-CH_2-O$$

$$MeO-CH_2-CH_2-O$$

61%

RX(170) OF 171 - 9 STEPS

н<sub>2</sub>с— с— сн 55%

CON: STEP(1) reflux STEP(2) reflux STEP(5) 160 deg C STEP(6) reflux STEP(7) reflux STEP(8) 60 deg C STEP(9) 100 deg C

- L3 ANSWER 6 OF 10 CASREACT COPYRIGHT 2008 ACS on STN
- 142:240390 Improved, high yield synthesis of 3H-quinazolin-4-ones, the key intermediates of recently developed drugs. Oerfi, Laszlo; Waczek, Frigyes; Pato, Janos; Varga, Istvan; Hegymegi-Barakonyi, Balint; Houghten, Richard A.; Keri, Gyoergy (Department of Pharmaceutical Chemistry, Semmelweis University, Budapest, Hung.). Current Medicinal Chemistry, 11(19), 2549-2553 (English) 2004. CODEN: CMCHE7. ISSN: 0929-8673. Publisher: Bentham Science Publishers Ltd..

61%

AB Purine bases and their bioisosteric analogs are widely used as building blocks in combinatorial chemical Recently a great number of fused pyrimidine derivs. became known as potential drug mols. against various types of proliferative diseases, caused by over-expression of protein kinases. One of the most important compound families are quinazolines: e.g. the best inhibitor of EGFR tyrosine kinase is PD153035 (6,7-dimethoxy-4-(3'-bromophenyl)amino-quinazoline) [2] and IRESSA (gefitinib, ZD1839) [3], developed from this compound family, is presently the only one approved and

granted drug by the FDA for the treatment of advanced non-small-cell lung cancer (NSCLD). KF31327 (3-sthyl-8-12-(4-hydroxymethyl-piperidino)benzylen ino]-2,3-dihydro-1H- imidazo[4,5-g]-quinazoline-2-thione dihydrochloride) from this group, showed significantly higher inhibitory activity on cyclic GMP-specific phosphodiesterase compared with those of sildenafil (Viagra). The synthetic procedures of the example compds. are based on imidoyl chloride intermediates that were prepared from the appropriate 3H-quinazoline-4-ones. Although the key intermediates, quinazoline-4-ones, have been known since more than hundred years, their synthetic procedures have been improved much only in the past ten years. In this paper we reviewed the efficient synthetic methods of quinazolin-4-ones; and presented a novel, reliable method for their synthesis. There was no considerable effect of microwave-, or traditional thermal activation on the yield and compound purity.

AcOH, HN:CHNH2, Formamide

NOTE: workup: hot aq. NaOH /charcoal decolorization and acidification (purity 87%)
CON: 2 hours, 160 deg C

L3 AMSWER 7 OF 10 CASREACT COPYRIGHT 2008 ACS on STN
142:219298 Process for preparation of 6,7-bis(2-methoxyethoxy)quinazolin-4one. Nishino, Shigeyoshi; Hirotsu, Kenji; Shima, Hidetaka; Oda, Hiroyuki;
Suzuki, Shinobu (Ube Industries, Ltd., Japan). PCT Int. Appl. WO
2005012264 Al 20050210, 18 pp. DeSIGNATED STATES: W: AE, AG, AL, AM, AT,
AU, AZ, BA, BB, BG, BR, BM, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK,
DM, DZ, EC, EE, EG, ES, FI, GB, GD, GB, GH, GM, HB, HU, DI, LI, IN, IS,
JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN,
MM, MX, MZ, NA, NI, NO, MZ, OM, PG, PH, PL, PT, RO, RU, SC, SD, SE, SG,
SK, SL, SY, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM,
ZM; RM: AT, BE, BF, BJ, CF, CG, CH, CI, CK, CY, DE, DK, ES, FI, FR, GA,
GB, GR, IE, IT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG, TR.
(Japanese). CODEN: PIXXD2. APPLICATION: WO 2004-JP10965 20040730.
PRIORITY: JP 2003-282696 20030730.

AB This invention pertains to a method for producing 6,7-bis(2-methoxy)quinazolin-4-one, which comprises reacting Et 2-amino-4,3-bis(2-methoxyethoxy) benzoate with a formic acid compound in the presence of an ammonium carboxylate. For example, Et 2-amino-4,5-bis(2-methoxyethoxy) benzoate (preparation given) was reacted with Me orthoformate in MeOH in the presence of NH4OAc to give the title compound (91%). This invention provides a convenient method to prepare the title compound from Et 2-amino-4,5-bis(2-methoxyethoxy) benzoate in high yield.

RX(4) OF 10

CON: 7 hours, 60 - 70 deg C

CON: STEP(1) 6 hours, 50 - 60 deg C STEP(2) 7 hours, 60 - 70 deg C RX(9) OF 10 - 3 STEPS

CON: STEP(1.1) room temperature -> 70 deg C; 2 hours, 70 deg C STEP(2) 6 hours, 50 - 60 deg C STEP(3) 7 hours, 60 - 70 deg C

RX(10) OF 10 - 4 STEPS

CON: STEP(1) 9 hours, 90 - 100 deg C
 STEP(2.1) room temperature -> 70 deg C; 2 hours, 70 deg C
 STEP(3) 6 hours, 50 - 60 deg C
 STEP(4) 7 hours, 60 - 70 deg C

L3 ANSWER 8 OF 10 CASREACT COPYRIGHT 2008 ACS on STN

139:164805 Process for producing 3,4-dihydroquinazolin-4-one derivatives.

Nishino, Shigeyoshi; Hirotsu, Kenji; Shima, Hidetaka; Harada, Takashi;
Oda, Hiroyuki; Takahashi, Takeshi; Suzuki, Shinobu (Ube Industries, Ltd.,
Japan). PCT Int. Appl. Wo 2003064399 Al 20030807, 101 pp. DESIGNATED

STATES: W: AE, AG, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, BZ, CA, CH,
CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM,
HR, HU, ID, IL, IN. IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU,
LV, MA, MD, MG, MK, MN, MW, MX, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SD,
SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA,
ZM, ZW; RW: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR,
GA, GB, GR, IE, IT, LU, MC, ML, MN, EN, LN, PT, SE, SN, TD, TG, TR.

(Japanese). CODEN: PIXXD2. APPLICATION: WO 2003-JP805 20030128. PRIORITY: JP 2002-17957 20020128; JP 2002-40929 20020219; JP 2002-82607 20020325; JP 2002-186443 20020610; JP 2002-178661 20020619; JP 2002-246657 20020827; JP 2002-336752 20021111: JP 2002-349456 20021201.

AB Disclosed is a process for producing a quinazolin-4-one derivative represented by the following formula (I) (wherein R1, R2, R3, and R4 each represents a group not participating in the reaction and R1, R2, R3, and R4 may be bonded to each other to form a ring) which comprises reacting an anthranilic acid derivative represented by the following formula (II) (wherein R1-R4 are defined as above; R5 represents hydrogen or a hydrocarbon group) with a formic acid derivative (in particular orthoformate ester) in the presence of an ammonium carboxylate. This process gives 3,4-dihydroquinazolin-4-one derivs. I, which are useful as intermediates for drugs and agrochems., from anthranilic acid derivs. in high yields in a simple method under mild conditions and is industrially suitable. Thus, 5-methoxy-4-(3-chloropropoxy)anthranilic acid Me ester 161.5, Me orthoformate 156.5, ammonium acetate 113.7 g, and 300 mL MeOH were added to a 1,000 mL stainless steel pressure vessel, allowed to react at 90-95° and 0.1-0.3 MPa for 8 h, treated with 600 mL H2O, stirred at 0-10° for 1 h, and filtered to give, after washing the crystals with 600 mL H2O and drying them at 60° under reduced pressure, 94% 6-methoxy-7-(3-chloropropoxy)quinazolin-4-one (152.8 g).

CH(OMe)3, NH4OAc, MeOH

NOTE: high pressure cyclocondensation in a stainless pressure vessel CON: 8 hours, 95 deg C

RX(12) OF 112

91%

NOTE: high pressure cyclocondensation in a stainless pressure vessel CON: 6 hours, 110 deg C

L3 ANSWER 9 OF 10 CASREACT COPYRIGHT 2008 ACS on STN

139:164804 Preparation of quinazolinone derivatives and 4,5-substituted-2-formylaminobenzamides as intermediates for said quinazolinones. Shirai, Masaahi; Furuya, Toshio (Ube Industries, Ltd., Japan). PCT Int. Appl. WO 2003064377 Al 20030807, 30 pp. DESIGNATED STATES: W: AE, AG, AL, AM, AT, AL, AZ, BA, BB, BG, BR, BY, BZ, CA, CH, CN, CO, CR, CU, CZ, DE, DK, DM, DZ, EC, EE, ES, FI, GB, GD, GE, GH, GM, RH, HU, ID, IL, IN, IS, JP, KE, KG, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MM, MZ, NO, NZ, OM, PH, PL, PT, RO, RU, SC, SD, SE, SG, SK, SL, TJ, TM, TN, TR, TT, TZ, UA, UG, US, UZ, VC, VN, YU, ZA, ZM, ZW; RN: AT, BE, BF, BJ, CF, CG, CH, CI, CM, CY, DE, DK, ES, FI, FR, GA, GB, GR, IE, ITT, LU, MC, ML, MR, NE, NL, PT, SE, SN, TD, TG, TR. (Japanese). CODEN: PIXXD2.

APPLICATION: WO 2003-JP562 20030122. PRIORITY: JP 2002-19583 20020129; JP 2002-77880 20020320.

G]

AB 4,5-Substituted-2-formylaminobenzamides (I) are produced by reacting the corresponding aminobenzamides with formic acid in an organic solvent, and I are further converted into 6,7-substituted-quinazolin-4-ones of the general formula II [R1, R2 = H, (un)substituted alkyl, etc.] by cyclization in the presence of a base. I are pharmaceutical intermediates. 6,7-Dimethoxy-3H-quinazolin-4-one was prepared in 93% yield by the above-mentioned process.

1. NaOH, Water 2. HC1, Water

MeO-CH<sub>2</sub>-CH<sub>2</sub>-O NH-CHO (step 1)

CON: STAGE(1) 30 minutes, 25 deg C, pH 13.5 STAGE(2) pH 7.6

RX(11) OF 12 - 2 STEPS

MeO-CHo-CHo-O

2.1. NaOH, Water 2.2. HCl, Water

MeO-CH<sub>2</sub>-CH<sub>2</sub>-O NH<sub>2</sub>

4%

- L3 ANSWER 10 OF 10 CASREACT COPYRIGHT 2008 ACS on STN
- 137:263058 Preparation of quinazoline derivative as intermediates for antitumor agent. Wang, Wi-Chi; Iseki, Elichi; Imamiya, Katsuyuki (Sumika Fine Chemicals Co., Ltd., Japan). Jpn. Kokai Tokkyo Koho JP 2002293773 A 20021009, 9 pp. (Japanese). CODEN: JKXXAF. APPLICATION: JP 2001-100183 20010330.
- AB 6,7-Bis(2-methoxyethoxy)-4(1H)-quinazolinone, an intermediate for the antitumor agent N-(3-ethylphenyl)-6,7-bis(2-methoxyethoxy)-4-quinazolinamine, is prepared in several steps from Et 3,4-dihydroxybenzoate and 2-methoxyethyl mesylate via cyclocondensation of Et 2-amino-4,5-bis(2-methoxyethoxy)benzoate with ammonium formate.

RX(5) OF 15

Ammonium formate, Formamide

RX(9) OF 15 - 2 STEPS

MeO-CH2-CH2-O

NO<sub>2</sub>

1. Pt, H2, MeOH

2. Ammonium formate, Formamide

RX(13) OF 15 - 3 STEPS

- 1. H2SO4, HNO3, AcOH 2. Pt, H2, MeOH 3. Ammonium formate, Formamide

$$\begin{array}{c} \text{MeO-CH}_2\text{--CH}_2\text{--O} \\ \text{MeO-CH}_2\text{--CH}_2\text{--O} \\ \end{array}$$

- 4. Ammonium formate, Formamide

80%

RX(15) OF 15 - 5 STEPS

- 1. MeCH2CH2OH, N-Methylmorpholine, THF
  - 2. Bu4N.I, K2CO3,
  - Me2CO 3. H2SO4, HNO3, AcOH 4. Pt, H2, MeOH
  - 5. Ammonium formate, Formamide